	Туре	L #	Hits	Search Text
1	BRS	L1	6405	pyrococcus horikoshi
2	BRS	L2	73472	pyrococcus (1n) horikoshi
3	BRS	L3	17	pyrococcus nearl horikoshi
4	BRS	L4	198	pyrococcus nearl horikoshii
5	BRS	L5	168	"15" and aminotransferase
6	BRS	L6	0	"15" near10 aminotransferase
7	BRS	L7	0	l4 near10 aminotransferase
8	BRS	T8	1	14 near20 aminotransferase

(FILE 'HOME' ENTERED AT 17:55:22 ON 10 AUG 2005)

FILE 'AGRICOLA, MEDLINE, CAPLUS, BIOSIS' ENTERED AT 17:55:25 ON 10 AUG 2005

663 S PYROCOCCUS (2N) HORIKOSHII L1

15 S L1 AND AMINOTRANSFERASE L2L3

9 DUP REM L2 (6 DUPLICATES REMOVED)

0 S L3 AND PY<1999 L4









. All Databases	PubMed	Nucieotida	Protein	Genome	Structure	OMM	PMC	Journals	Br
Search PubMed		for 2.6.	1.57[EC/R				PIC	view Go	C
•	/.	w			384		*		
	₩ Lir	nits Previe	w/Index	History	Clipboard	Details			

## Limits: Publication Date to 1998

About Entrez

Text Version

Entrez PubMed Overview Help | FAQ Tutorial New/Noteworthy E-Utilities

PubMed Services
Journals Database
MeSH Database
Single Citation Matcher
Batch Citation Matcher
Clinical Queries
Special Queries
LinkOut
My NCBI (Cubby)

Related Resources
Order Documents
NLM Catalog
NLM Gateway
TOXNET
Consumer Health
Clinical Alerts
ClinicalTrials.gov
PubMed Central

• Search History will be lost after eight hours of inactivity.

- To combine searches use # before search number, e.g., #2 AND #6.
- Search numbers may not be continuous; all searches are represented.
- Click on query # to add to strategy

Search	Most Recent Queries	Time	Result
<u>#26</u>	Search 2.6.1.57[EC/RN Number] Field: All Fields, Limits: Publication Date to 1998	17:54:16	<u>29</u>
<u>#25</u>	Search 2.6.1.57[EC/RN Number] Limits: ignored	17:54:09	<u>37</u>
<u>#23</u>	Search Pyrococcus horikoshii Limits: Publication Date to 1998	17:52:49	<u>7</u>
#21	Search kawarabayasi pyrococcus Field: All Fields, Limits: Publication Date to 1998	17:52:18	<u>3</u>
<u>#20</u>	Search kawarabayasi pyrococcus	17:52:10	<u>20</u>
<u>#19</u>	Search kawarabayasi	17:52:03	<u>69</u>
<u>#13</u>	Search pyrococcus horikoshii aminotransferase	17:51:32	<u>3</u>
<u>#18</u>	Related Articles for PubMed (Select 10671523)	17:50:42	<u> 191</u>
#17	Search pyrococcus horikoshii aminotransferase kawarabayasi	17:50:33	1
<u>#12</u>	Search pyrococcus horikoshii	17:48:27	<u>161</u>
<u>#11</u>	Search pyrococcus horikoshi	17:48:21	<u>2</u>
<u>#10</u>	Search kurabayashi tanaka mercury	16:14:08	<u>1</u>
<u>#9</u>	Search kurabayashi tanaka	16:14:01	<u>88</u>
<u>#8</u>	Search carroll minton	16:13:34	<u>3</u>
<u>#6</u>	Search carroll and minton	14:46:37	<u>3</u>
<u>#5</u>	Search (deinococcus [ti] OR radiodurans [ti])	14:23:00	<u>326</u>
<u>#4</u>	Search deinococcus bioremediation	14:21:16	7
<u>#3</u>	Search deinococcus radiodurans bioremediation	14:21:04	<u>7</u>
<u>#1</u>	Search ferreira deinococcus	14:20:31	· <u>1</u>

**Clear History** 

Write to the Help Desk
NCBI | NLM | NIH
Department of Health & Human Services
Privacy Statement | Freedom of Information Act | Disclaimer

Jul 26 2005 04:43:15

cited previously

```
RESULT 1
059096
ID
    059096
                PRELIMINARY;
                                  PRT;
                                         389 AA.
AC
    059096;
DT
     01-AUG-1998 (TrEMBLrel. 07, Created)
DT
     01-AUG-1998 (TrEMBLrel. 07, Last sequence update)
DT
     01-0CT-2003 (TrEMBLrel. 25, Last annotation update)
DΕ
     389aa long hypothetical aspartate aminotransferase.
GN
    OrderedLocusNames=PH1371;
os
    Pyrococcus horikoshii.
OC
    Archaea; Euryarchaeota; Thermococci; Thermococcales; Thermococcaceae;
OC
     Pyrococcus.
XO,
    NCBI TaxID=53953;
RN ·
     [1]
RP
     SEQUENCE FROM N.A.
RC
     STRAIN=OT3;
RX
    MEDLINE=98344137; PubMed=9679194;
RA
    Kawarabayasi Y., Sawada M., Horikawa H., Haikawa Y., Hino Y.,
RA
    Yamamoto S., Sekine M., Baba S.-I., Kosugi H., Hosoyama A., Nagai Y.,
RA
     Sakai M., Ogura K., Otsuka R., Nakazawa H., Takamiya M., Ohfuku Y.,
RA
     Funahashi T., Tanaka T., Kudoh Y., Yamazaki J., Kushida N., Oguchi A.,
RA
    Aoki K.-I., Yoshizawa T., Nakamura Y., Robb F.T., Horikoshi K.,
RA
     Masuchi Y., Shizuya H., Kikuchi H.;
RT
     "Complete sequence and gene organization of the genome of a hyper-
RT
     thermophilic archaebacterium, Pyrococcus horikoshii OT3.";
RL
    DNA Res. 5:55-76(1998).
CC
     -!- COFACTOR: Pyridoxal phosphate (By similarity).
CC
     -!- SIMILARITY: Belongs to the class-I pyridoxal-phosphate-dependent
        aminotransferase family.
CC
DR
    EMBL; AP000006; BAA30477.1; -.
DR
    PIR; E71009; E71009.
DR
    PDB; 1DJU; X-ray; A/B=2-389.
DR
     PDB; 1GD9; X-ray; A/B=1-389.
DR
     PDB; 1GDE; X-ray; A/B=1-389.
DR
     GO; GO:0016847; F:1-aminocyclopropane-1-carboxylate synthase . . .; IEA.
DR
    GO; GO:0008483; F:transaminase activity; IEA.
DR
    GO; GO:0009058; P:biosynthesis; IEA.
DR
     InterPro; IPR001176; ACC_synthase.
DR
     InterPro; IPR004839; Aminotrans I/II.
DR
     InterPro; IPR004838; NHtransf 1 BS.
DR
    Pfam; PF00155; Aminotran 1 2; 1.
DR
     PRINTS; PR00753; ACCSYNTHASE.
DR
    PROSITE; PS00105; AA_TRANSFER_CLASS 1; 1.
KW
    Aminotransferase; Complete proteome; Hypothetical protein;
KW
     Pyridoxal phosphate; Transferase.
SQ
    SEQUENCE
               389 AA; 43900 MW; F2A3E919F0C2A7FF CRC64;
  Query Match
                         100.0%; Score 1985; DB 2;
                                                     Length 389;
  Best Local Similarity
                         100.0%; Pred. No. 1e-130;
  Matches 389; Conservative
                                0; Mismatches
                                                     Indels.
Qy
           1 MALSDRLELVSASEIRKLFDIAAGMKDVISLGIGEPDFDTPOHIKEYAKEALDKGLTHYG 60
             Db
           1 MALSDRLELVSASEIRKLFDIAAGMKDVISLGIGEPDFDTPQHIKEYAKEALDKGLTHYG 60
          61 PNIGLLELREAIAEKLKKQNGIEADPKTEIMVLLGANQAFLMGLSAFLKDGEEVLIPTPA 120
Qу
```

٠.				
	Db	61	PNIGLLELREAIAEKLKKQNGIEADPKTEIMVLLGANQAFLMGLSAFLKDGEEVLIPTPA	120
	Qy	121	FVSYAPAVILAGGKPVEVPTYEEDEFRLNVDBLKKYVTDKTRALIINSPCNPTGAVLTKK	180
	Db	121		180
	Qy	181	DLEEIADFVVEHDLIVISDEVYEHFIYDDARHYSIASLDGMFERTITVNGFSKTFAMTGW	240
	Db	181	DLEEIADFVVEHDLIVISDEVYEHFIYDDARHYSIASLDGMFERTITVNGFSKTFAMTGW	240
	Qy ·	241	RLGFVAAPSWIIERMVKFQMYNATCPVTFIQYAAAKALKDERSWKAVEEMRKEYDRRRKL	300
	Db	241	RLGFVAAPSWIIERMVKFQMYNATCPVTFIQYAAAKALKDERSWKAVEEMRKEYDRRRKL	300
	Qy	301	VWKRLNEMGLPTVKPKGAFYIFPRIRDTGLTSKKFSELMLKEARVAVVPGSAFGKAGEGY	360
	Db	301	VWKRLNEMGLPTVKPKGAFYIFPRIRDTGLTSKKFSELMLKEARVAVVPGSAFGKAGEGY	360
	Qу	361	VRISYATAYEKLEEAMDRMERVLKERKLV 389	
	Db	361	VRISYATAYEKLEEAMDRMERVLKERKLV 389	
				•
				·

·

```
E71009
probable aromatic-amino-acid transaminase (EC 2.6.1.57) PH1371 [similarity] -
Pyrococcus horikoshii
C; Species: Pyrococcus horikoshii
C;Date: 14-Aug-1998 #sequence revision 14-Aug-1998 #text change 12-Jul-2004
C; Accession: E71009
R; Kawarabayasi, Y.; Sawada, M.; Horikawa, H.; Haikawa, Y.; Hino, Y.; Yamamoto,
S.; Sekine, M.; Baba, S.; Kosugi, H.; Hosoyama, A.; Nagai, Y.; Sakai, M.; Ogura,
K.; Otsuka, R.; Nakazawa, H.; Takamiya, M.; Ohfuku, Y.; Funahashi, T.; Tanaka,
T.; Kudoh, Y.; Yamazaki, J.; Kushida, N.; Oquchi, A.; Aoki, K.; Yoshizawa, T.;
Nakamura, Y.; Robb, F.T.; Horikoshi, K.; Masuchi, Y.; Shizuya, H.; Kikuchi, H.
DNA Res. 5, 55-76, 1998
A; Title: Complete sequence and gene organization of the genome of a hyper-
thermophilic archaebacterium, Pyrococcus horikoshii OT3.
A; Reference number: A71000; MUID: 98344137; PMID: 9679194
A: Accession: E71009
A; Status: preliminary; nucleic acid sequence not shown; translation not shown
A; Molecule type: DNA
A; Residues: 1-389 < KAW>
A; Cross-references: UNIPROT:059096; GB:AP000006; NID:g3236133; PIDN:BAA30477.1;
PID:g3257794
A; Experimental source: strain OT3
A; Note: this accession replaces an interim accession for a sequence replaced by
GenBank
C:Genetics:
A; Gene: PH1371
C; Keywords: aminotransferase; phosphoprotein; pyridoxal phosphate
F;233/Binding site: pyridoxal phosphate (Lys) (covalent) #status predicted
 Query Match
                       100.0%; Score 1985; DB 2; Length 389;
 Best Local Similarity
                      100.0%; Pred. No. 4.4e-135;
 Matches 389; Conservative
                            0: Mismatches
                                             0: Indels
                                                            Gaps
                                                         0:
          1 MALSDRLELVSASEIRKLFDIAAGMKDVISLGIGEPDFDTPQHIKEYAKEALDKGLTHYG 60
Qy
            Db
          1 MALSDRLELVSASEIRKLFDIAAGMKDVISLGIGEPDFDTPOHIKEYAKEALDKGLTHYG 60
Qy
         61 PNIGLLELREAIAEKLKKONGIEADPKTEIMVLLGANOAFLMGLŠAFLKDGEEVLIPTPA 120
            Db
         61 PNIGLLELREAIAEKLKKONGIEADPKTEIMVLLGANQAFLMGLSAFLKDGEEVLIPTPA 120
        121 FVSYAPAVILAGGKPVEVPTYEEDEFRLNVDELKKYVTDKTRALIINSPCNPTGAVLTKK 180
            Db
        121 FVSYAPAVILAGGKPVEVPTYEEDEFRLNVDELKKYVTDKTRALIINSPCNPTGAVLTKK 180
Qу
        181 DLEEIADFVVEHDLIVISDEVYEHFIYDDARHYSIASLDGMFERTITVNGFSKTFAMTGW 240
            Db
        181 DLBEIADFVVEHDLIVISDEVYEHFIYDDARHYSIASLDGMFERTITVNGFSKTFAMTGW 240
Qу
        241 RLGFVAAPSWIIERMVKFOMYNATCPVTFIOYAAAKALKDERSWKAVEEMRKEYDRRRKL 300
            Db
        241 RLGFVAAPSWIIERMVKFQMYNATCPVTFIQYAAAKALKDERSWKAVEEMRKEYDRRRKL 300
        301 VWKRLNEMGLPTVKPKGAFYIFPRIRDTGLTSKKFSELMLKEARVAVVPGSAFGKAGEGY 360
Qу
            301 VWKRLNEMGLPTVKPKGAFYIFPRIRDTGLTSKKFSELMLKBARVAVVPGSAFGKAGEGY 360
Db
```

RESULT 1

Qy	361 VRISYATAYEKLEEAMDRMERVLKERKLV 3	389
Db	361 VRISYATAYEKLEEAMDRMERVLKERKLV 3	389

:

.

•